EXHIBIT 3

```
EMBL; AE004969; AAW89056.1; -; Genomic DNA.
DR
    RefSeg; YP 207468.1; -.
DR
   HSSP; P13000; 1BYI.
DR
    STRING; Q5F9T1; -.
DR
    GeneID; 3281703; -.
DR
   GenomeReviews; AE004969 GR; NG00309.
DR
   KEGG; ngo:NGO0309; -.
DR
   NMPDR; fig|242231.4.peg.499; -.
DR
   HOGENOM; HBG650065; -.
DR
   OMA; LLTIDYI; -.
DR
    BioCvc; NGON242231:NGO0309-MON; -.
    GO; GO:0005524; F:ATP binding; IEA:HAMAP.
DR
    GO; GO:0004141; F:dethiobiotin synthase activity; IEA:HAMAP.
DR
    GO; GO:0000287; F:magnesium ion binding; IEA:HAMAP.
    GO: GO:0009102; P:biotin biosynthetic process; IEA:HAMAP.
DR
   HAMAP; MF 00336; -; 1.
    InterPro; IPR004472; BioD synth.
DR
DR
    PIRSF; PIRSF006755; DTB synth; 1.
PE
    3: Inferred from homology;
KW
   ATP-binding; Biotin biosynthesis; Complete proteome; Ligase;
KW
   Magnesium; Nucleotide-binding.
FT
   CHAIN
                       215
                                 Dethiobiotin synthetase.
FT
                                 /FTId=PRO 0000302532.
FT
                       17
                                 ATP (By similarity).
   NP BIND
    SEQUENCE 215 AA; 23676 MW; 48CB57B353B7CD95 CRC64;
    MKGVYFVSGI DTDIGKTVAT GMLAKQLLQQ GKSVITQKPV QTGCQDIAED IAVHRKIMGI
    PMQEADEQRL TMPEIFSHPA SPHLAARLDG RGLDLDKIRT ATQELAAQYE VVLVEGAGGL
    MVPLTEKLLT IDHIQQQAYP VILVTSGRLG SINHTLLSFV VLKQYGIRLH SLVFNHIHDS
    RDAHVAODSL NYLOCRLKAD FPEAEWMELA KTGAV
    EMBL; AE004969; AAW90296.1; ALT INIT; Genomic DNA.
DR
    RefSeq; YP 208708.1; -.
DR
    STRING; 05F691; -.
DR
    GeneID: 3281272: -.
DR
    GenomeReviews; AE004969 GR; NG01671.
DR
   KEGG; ngo:NGO1671; -.
DR
   NMPDR; fig|242231.4.peg.1812; -.
DR
    HOGENOM; HBG650290; -.
DR
   OMA; IIAREVV; -.
DR
   BioCvc; NGON242231:NGO1671-MON; -.
DR
    GO; GO:0005737; C:cytoplasm; IEA:UniProtKB-SubCell.
DR
    GO; GO:0005524; F:ATP binding; IEA:HAMAP.
DR
    GO; GO:0004140; F:dephospho-CoA kinase activity; IEA:HAMAP.
DR
    GO; GO:0015937; P:coenzyme A biosynthetic process; IEA:HAMAP.
DR
    HAMAP; MF 00376; -; 1.
    InterPro; IPR001977; Depp CoAkinase.
DR
DR
   PROSITE; PS51219; DPCK; 1.
PΕ
    3: Inferred from homology:
   ATP-binding; Coenzyme A biosynthesis; Complete proteome;
Cvtoplasm:
KW
    Kinase; Nucleotide-binding; Transferase.
    CHAIN
                  1 210
                                 Dephospho-CoA kinase.
```

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FT
                                 /FTId=PRO 0000243307.
FT
    DOMAIN
                       202
                  4
                                 DPCK.
FT
    NP BIND
                  9
                       16
                                 ATP (Potential).
SO
    SEQUENCE 210 AA; 22950 MW; 1443ED0A3D4057B5 CRC64;
    MTAWVGLTGG IGSGKSAAAO YFADLGVPRI DADAAAHSLT ASDGIALPEI RRLFGDTVFD
    TOGLLRRDIL RKEIFASPSR KALLESVMLP LIFSEIKKOO ETFTDAVYGI VEIPLLTEKR
    OFISLIRRVL TISAPLEKRI GRVMARSGLT RGEVADIISH OASESERLLL ADDVLLNDGS
    LKSLREKTML LHAFYSGIFA SKPTOGKHNG
    EMBL; AE004969; AAW88861.1; ALT INIT; Genomic DNA.
DR
    RefSeq; YP 207273.1; -.
DR
    HSSP; POA6P7; 1PUI.
DR
    STRING; Q5FAC6; -.
DR
    GeneID; 3282443; -.
DR
    GenomeReviews; AE004969 GR; NG00100.
DR
    KEGG; ngo:NGO0100; -.
DR
   NMPDR; fig|242231.4.peg.842; -.
DR
   HOGENOM: HBG447097: -.
DR
   OMA; THFVTSA; -.
DR
    BioCyc; NGON242231:NGO0100-MON; -.
DR
    GO; GO:0005622; C:intracellular; IEA:InterPro.
DR
    GO; GO:0005525; F:GTP binding; IEA:HAMAP.
DR
    GO; GO:0000917; P:barrier septum formation; IEA:HAMAP.
DR
    GO; GO:0007049; P:cell cycle; IEA:UniProtKB-KW.
   HAMAP; MF 00321; -; 1.
DR
DR
    InterPro; IPR019987; GTP-bd ribosome bio YsxC.
    InterPro; IPR002917; MMR HSR1 GTP bd.
DR
DR
    Pfam; PF01926; MMR HSR1; 1.
PE.
    3: Inferred from homology;
KW
   Cell cycle; Cell division; Complete proteome; GTP-binding;
KW
    Nucleotide-binding; Septation.
FT
   CHAIN
                  1
                       209
                                 Probable GTP-binding protein engB.
FT
                                 /FTId=PRO 0000266903.
FT
    NP BIND
                 30
                        37
                                 GTP (Potential).
FT
    NP BIND
                 75
                        79
                                 GTP (Potential).
FT
    NP BIND
                142
                       145
                                 GTP (Potential).
    SEQUENCE
               209 AA; 23600 MW; 79742D484A0A1ABF CRC64;
SQ
    MNLFONAKFF TTVNHLKDLP DTPLEIAFVG RSNAGKSSAI NTLTNHVRLA YVSKTPGRTO
    HINFFELONG NFMVDLPGYG YAQVPEAVRA HWVNLLGDYL RHRKQLIGLV LIMDARHPLK
    ELDIRMLDFF HTTGRPVHIL LSKADKLSKN EOIKTLSOVK KLLKPYSDRO NISVOLFSSL
    KKQGIDEANR TVGSWFDAAD AAASSPEEN
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RefSeq; YP 208309.1; -.
    STRING; Q5F7E0; -.
DR
DR
   GeneID; 3282595; -.
DR
   GenomeReviews; AE004969 GR; NG01238.
DR
   KEGG; ngo:NG01238; -.
DR
   NMPDR; fig|242231.4.peg.1300; -.
DR
   HOGENOM: HBG391868: -.
DR OMA; LLMFRAE; -.
DR
   BioCvc; NGON242231:NGO1238-MON; -.
DR
    GO; GO:0005737; C:cvtoplasm; IEA:UniProtKB-SubCell.
    GO; GO:0005524; F:ATP binding; IEA:UniProtKB-KW.
DR
    GO; GO:0003879; F:ATP phosphoribosyltransferase activity;
IEA: HAMAP.
    GO; GO:0000105; P:histidine biosynthetic process; IEA:HAMAP.
DR
    HAMAP; MF 01018; -; 1.
    InterPro; IPR001348; ATP PRibTrfase.
DR
    InterPro; IPR013820; ATP PRibTrfase cat.
DR
   InterPro; IPR018198; ATP PRibTrfase CS.
DR
   PANTHER; PTHR21403; ATP phspho trans; 1.
DR
   Pfam; PF01634; HisG; 1.
DR PROSITE; PS01316; ATP P PHORIBOSYLTR; 1.
PE
    3: Inferred from homology;
KW
    Amino-acid biosynthesis; ATP-binding; Complete proteome;
Cytoplasm;
KW
    Glycosyltransferase; Histidine biosynthesis; Nucleotide-binding;
KW
    Transferase.
FT
    CHAIN
                        221
                                  ATP phosphoribosyltransferase.
FT
                                  /FTId=PRO 0000229320.
SQ
    SEQUENCE 221 AA; 23776 MW; 74D2248667173BD0 CRC64;
    MODNALTIAL SKGRIFEETL PLLAAAGIAP TEEPEKSRKL IIGTNHENIR LVIVRATDVP
    TYVRYGAADF GIAGKDVLIE HGGTGLYRPL DLEIAKCRMM VAVRKGFDYE AASQPGCRLK
     IATKYPEIAA SHFAGKGVHV DIIKLYGSME LAPLVGLSDA IVDLVSTGNT LKANGLEAVE
     HIVDISSYLV VNKAALKTKY ALLEPIIQSF GGAVKAKWAF I
DR
    EMBL; AE004969; AAW88966.1; -; Genomic DNA.
DR
    RefSeq; YP 207378.1; -.
DR
    STRING; Q5FA21; -.
DR
    GeneID; 3281167; -.
DR
    GenomeReviews; AE004969 GR; NG00213.
DR
    KEGG: ngo:NGO0213: -.
   NMPDR; fig|242231.4.peg.252; -.
DR
DR
   HOGENOM; HBG292341; -.
DR
    OMA; SVRFAFE; -.
DR
    BioCyc; NGON242231:NGO0213-MON; -.
DR
    GO; GO:0005737; C:cytoplasm; IEA:UniProtKB-SubCell.
    GO; GO:0000107; F:imidazoleglycerol-phosphate synthase activity;
IEA: HAMAP.
DR
    GO; GO:0006541; P:glutamine metabolic process; IEA:UniProtKB-KW.
DR
    GO; GO:0000105; P:histidine biosynthetic process; IEA:HAMAP.
DR
    HAMAP; MF_00278; -; 1.
InterPro; IPR017926; GATASE_1.
DR
DR
   InterPro; IPR000991; GATase class1 C.
DR
   InterPro; IPR010139; Imidazole-glycPsynth HisH.
DR InterPro; IPR016226; Imidazole-GPS HisH.
DR
   Pfam; PF00117; GATase; 1.
```

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PIRSF; PIRSF000495; Amidotransf hisH; 1.
    PROSITE; PS51273; GATASE TYPE 1; 1.
DR
PE
    3: Inferred from homology;
KW
    Amino-acid biosynthesis; Complete proteome; Cytoplasm;
KW
    Glutamine amidotransferase; Histidine biosynthesis; Transferase.
FΤ
    CHAIN
                  1
                       212
                                 Imidazole glycerol phosphate synthase
FT
                                 subunit hisH.
FT
                                 /FTId=PRO 0000231736.
FT
    DOMAIN
                       212
                                 Glutamine amidotransferase type-1.
FT
    ACT SITE
                 85
                        85
                                 Nucleophile (By similarity).
FT
    ACT_SITE
                194
                       194
                                 By similarity.
FT
    ACT SITE
               196
                       196
                                 By similarity.
SO
    SEQUENCE
              212 AA; 23570 MW; BDDFE90E1A43ED5A CRC64;
    MOTALIDYGM GNLHSVLKSV RTAGOLAGKN TKIFLSGDPD RVSRADKVIF PGOGAMPDCM
    AALTRGGLDE AVKDALKNKP FFGICVGAQL LFDHSEEGNT DGLGWFGGKV RRFARDLRDP
    QGCRLKVPHM GWNTVRQTQN HPLFQGIPQN TRFYFVHSYY FAPENPETIL GESDYPSPFA
    CIVGKDNVFA TOFHTEKSHD AGLTMLKNFL NW
    EMBL; AE004969; AAW90546.1; -; Genomic DNA.
DR
    RefSeg; YP 208958.1; -.
   STRING; Q5F5J1; -.
DR
DR
   GeneID; 3282686; -.
    GenomeReviews: AE004969 GR: NG01933.
DR
DR
    KEGG; ngo:NGO1933; -.
   NMPDR; fig|242231.4.peg.1749; -.
DR
DR
    HOGENOM; O5F5J1; -.
DR
    OMA; GACYHDL; -.
DR
    PhylomeDB; O5F5J1; -.
DR
    BioCyc; NGON242231:NGO1933-MON; -.
DR
    GO; GO:0008936; F:nicotinamidase activity; IEA:EC.
DR
   GO; GO:0008152; P:metabolic process; IEA:InterPro.
    InterPro: IPR000868; Isochorismatase-like.
DR
DR
    Gene3D; G3DSA:3.40.50.850; Isochorismatase hydro; 1.
DR
    Pfam: PF00857: Isochorismatase: 1.
PE
    4: Predicted;
KW
    Complete proteome: Hydrolase.
    SEQUENCE 211 AA; 23439 MW; D1EFDE053D35A3EC CRC64;
SO
    MIVSIDVDAQ KTFTPLCPDE LPVNEGHLIV EELNAQAALA DLRVMTKDAH HMAAKWLVDN
    PVDMLKPTGF SDADLTWVAH AMVGTRGYEL LDGLPSVKEY DYCVWKGVDP ELHPYGACFH
    DIEEKLSTGL IEWLRCONTN MVIVGGLATD YCVKTTVLQL LKGGRWQVIV NEAACRGIAP
    DTIEAAWOEM RSSGAIILKN AEKIKKYINN O
    EMBL; AE004969; AAW90184.1; -; Genomic DNA.
DR
   RefSeg; YP 208596.1; -.
DR
    STRING; Q5F6K3; -.
DR
    GeneID; 3281487; -.
    GenomeReviews; AE004969 GR; NG01548.
DR
DR
    KEGG; ngo:NGO1548; -.
DR
   NMPDR: fig|242231.4.peg.1610; -.
DR
   HOGENOM; O5F6K3; -.
DR
   OMA; YERIAQP; -.
DR
   PhylomeDB; O5F6K3; -.
   BioCyc; NGON242231:NGO1548-MON; -.
```

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GO; GO:0030288; C:outer membrane-bounded periplasmic space;
IEA: InterPro.
    GO; GO:0003756; F:protein disulfide isomerase activity; IEA:EC.
    GO; GO:0015035; F:protein disulfide oxidoreductase activity;
IEA: InterPro.
   GO; GO:0045454; P:cell redox homeostasis; IEA:InterPro.
   InterPro: IPR001853; OxRdtase DSBA.
   InterPro; IPR017936; Thioredoxin-like.
DR
   InterPro; IPR012336; Thioredoxin-like fold.
DR
    InterPro; IPR017937; Thioredoxin CS.
    InterPro; IPR012335; Thioredoxin fold.
   Gene3D; G3DSA:3.40.30.10; Thioredoxin fold; 1.
   Pfam; PF01323; DSBA; 1.
   PROSITE; PS00194; THIOREDOXIN 1; 1.
DR
   PROSITE; PS51352; THIOREDOXIN 2; 1.
PE
    4: Predicted:
KW
    Complete proteome; Isomerase.
    SEQUENCE
              214 AA; 24003 MW; 31A85BBA5EB3CE18 CRC64;
    MKFKHLLPLL LSAVLSAOAY ALTEGEDYLV LDKPIPOEOP GKIEVLEFFG YFCVHCHHFD
    PLLLKLGKAL PSDTYLRTEH VVWRPEMLGL ARMAAAVKLS GLKYQANSAV FKAVYEQKIR
    LENRAVAGKW ALSOKGFDGK KLMRAYDSPE AAAVALKMOK LTEOYGIDST PTVIVGGKYR
    VIFNNGFDGG VHTIKELVAK VREERKRQTP AVQK
DR
    EMBL; AE004969; AAW90118.1; -; Genomic DNA.
    RefSeg; YP 208530.1; -.
DR
DR
   STRING; Q5F6R9; -.
DR GeneID; 3281613; -.
DR GenomeReviews; AE004969 GR; NG01479.
DR
   KEGG; ngo:NGO1479; -.
DR
    NMPDR; fig|242231.4.peg.1454; -.
DR
    HOGENOM; Q5F6R9; -.
DR
    PhylomeDB; Q5F6R9; -.
DR
   BioCyc; NGON242231:NGO1479-MON; -.
    4: Predicted;
KW
    Complete proteome.
SO
    SEQUENCE 209 AA; 23458 MW; B72DD3AA633EB0D3 CRC64;
    MEVOLPKIKT VRVMLAGMTA QQESVFKMAF KMHNTTRYET VSPSDGSAVP DLVLADTDAE
    GGFELWKELA GRYKAIPVAV CSEKVPDSEV PYLPKPIRFE TLFPMLRKLL QGENVYGKSF
    IAPADRSAKN NGNVORTVTI ROFNPNKGLL GALRFAEKNR ODIAILHGNK PVLNCFPLDT
    TGFADRKCVK TRRIVORRKF AGOLODCSR
    EMBL; AE004969; AAW90087.1; -; Genomic DNA.
DR
DR
    RefSeg; YP 208499.1; -.
    STRING: O5F6V0: -.
DR
    GeneID; 3281693; -.
DR
   GenomeReviews; AE004969 GR; NG01445.
DR
    KEGG; ngo:NGO1445; -.
DR
    NMPDR; fig|242231.4.peg.1423; -.
DR
   HOGENOM; O5F6V0; -.
DR
   OMA; ETRANAY; -.
DR PhvlomeDB; O5F6V0; -.
DR BioCyc; NGON242231:NGO1445-MON; -.
DR GO; GO:0019867; C:outer membrane; IEA:InterPro.
   GO; GO:0009405; P:pathogenesis; IEA:InterPro.
DR
    InterPro; IPR008640; Hep Hag.
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- DR Pfam; PF05658; Hep Hag; 4.
- PE 4: Predicted;
- KW Complete proteome.
- SQ SEQUENCE 222 AA; 22826 MW; C024A19C5373E774 CRC64;

MGLRQKLRGI IPNLATSIGT SAEANAPGAL ALGGSSEASK KFSIAEGYLA SSDGYGAIAI GSAAKIKOLE KGTINHIVCN DHKGLYVDAD GNVTKITVRT ESEKDILGRY GOTYGAVALG FRSSSHNLFA SSFGAFSTAT ALESLAVGDS SQSTGYRSAT FGSHSRALAE ESLALGYETR ANAYGSVALG AESVAMEENT VSVSSDTLKR KINNVADGTE DL

- DR EMBL; AE004969; AAW90078.1; -; Genomic DNA.
- DR RefSeg; YP 208490.1; -.
- DR GeneID; 3281702; -.
- DR GenomeReviews; AE004969 GR; NG01435.
- DR KEGG; ngo:NGO1435; -.
- DR NMPDR; fig|242231.4.peg.767; -.
- DR HOGENOM; O5F6V9; -.
- DR OMA; FFARANT; -.
- DR PhylomeDB; O5F6V9; -.
- DR BioCyc; NGON242231:NGO1435-MON; -.
- DR InterPro; IPR004676; Cd-R transporter.
- DR InterPro; IPR018388; Cd-R_transporter_subgr.
- DR Pfam; PF03596; Cad; 1.
- DR TIGRFAMs; TIGR00779; cad; 1.
- PE 4: Predicted:
- KW Complete proteome.
- SQ SEQUENCE 208 AA; 23177 MW; 1297237EA5DD043C CRC64;

MRCFMFSTVI TAAVLYIATA VDLLVILLIF FARANTRKEY RDIYIGQYLG SVILILVSLF LAPVLNYVYB KWIGLIGLI PIYLGIKVAI YDDCBGEKRA KKELDEKGLS KLVGIVALDY VASCGADDIG LFVPYEVTLD LVDLLVYLLV FLILIFVLVY TAQRLANISG VGEIVEKFSR WIMAVIYIGL GLFIIIENNT IRTIISII